Electronic spin transition of iron in the Earth's lower mantle

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Abstract Silicate perovskite and ferropericlase are thought to be the primary constituents of the lower mantle, whereas silicate post-perovskite is more likely found in the lowermost mantle. Because these minerals contain certain amounts of iron, their properties and, consequently, those of the deep mantle, are strongly influenced by iron's spin and valence states. A high-spin to low-spin crossover in ferropericlase has been observed to occur in the middle part of the lower-mantle conditions. Recent Mössbauer results consistently show that Fe^{2+} predominantly exhibits extremely high quadrupole splittings in perovskite and post-perovskite, whereas a high-spin to low-spin transition of Fe^{3+} in the octahedral site occurs at high pressures. These results provide a new venue for discussion of the effects of the spin and valence states of iron on the physical and chemical properties of the lower mantle.

Keywords Spin transition • Lower mantle • Mössbauer spectroscopy • Diamond anvil cell

1 Introduction

Earth's lower mantle, the most voluminous layer of the planet, is subjected to extreme pressure-temperature (P-T) conditions of 23–136 GPa and approximately 1,600–3,500 K. The lower mantle, as described in terms of a pyrolitic compositional model, is mainly comprised of ferropericlase ((Mg,Fe)O; Fp; \sim 33% by volume), aluminous silicate perovskite (Al-(Mg,Fe)SiO₃; \sim 62% by volume; hereafter called perovskite or Pv), and \sim 5% calcium silicate Pv (CaSiO₃, abbreviated CaPv) [1] (Figs. 1 and 2). Recent studies show that Pv transforms into silicate post-perovskite (PPv) within the D" layer, just above the core-mantle region. The implications of

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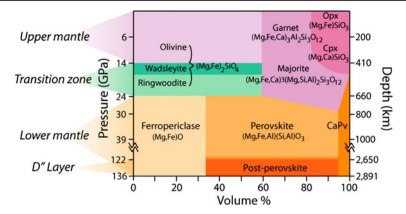


Fig. 1 Approximate mineral distribution within Earth's interior as a function of depth, pressure and volume%. Mineral abbreviation: Opx and cpx, orthopyroxene and clinopyroxene, respectively; CaPv, calcium silicate perovskite

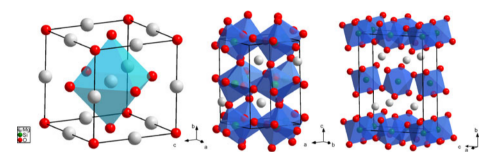


Fig. 2 Crystal structures of lower-mantle ferropericlase (*left*), silicate perovskite (*middle*), and silicate post-perovskite (*right*). Cubic ferropericlase is in the rock salt structure ($Fm\bar{3}m$); orthorhombic perovskite is in the *Pbnm* space group; orthorhombic post-perovskite is in the *Cmcm* space group

these minerals' physical and chemical properties are needed to explain the seismic, geochemical, geodynamic, and geomagnetic observations of the lower mantle. As the most abundant 3d transition metal in Earth's interior, iron affects many of the chemical and physical properties of the lower-mantle minerals [1]. Amongst the majority of these, iron exists mainly as ferrous iron (Fe²⁺), with approximately 20% concentrated in Fp, whereas Pv and PPv, with approximately 10% total iron, can contain very high amounts of ferric iron (Fe³⁺) [1].

Due to its partially filled 3*d*-electron orbitals, iron's electronic valence and spin states in the host phases can cause a dynamic range of physical and chemical properties of the lower mantle [1]. Previous studies have shown that Fe spin transitions and variations in Fe valence states can cause changes in density, elastic properties, electrical conductivity and radiative thermal conductivity of the lowermantle minerals. Here, literature data on the spin and valence states of iron in Fp, Pv, and PPv at relevant *P-T* conditions have been evaluated. Focus is placed on the use of the hyperfine parameters, quadrupole splittings (QS) and chemical shifts (CS), seen in Mössbauer spectroscopic analyses including synchrotron Mössbauer



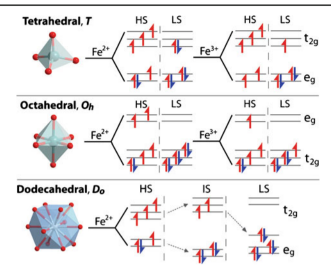


Fig. 3 Crystal field splitting diagrams for iron in tetrahedral, octahedral, and dodecahedral sites in the lower-mantle minerals. Iron is shown as a (2+) or (3+) cation in high-spin, intermediate-spin (dodecahedral site), and low-spin electronic configurations. The crystal field splitting energy (CFSE) can be altered by pressure, temperature, and/or composition. The energy of the e_g orbitals (those which are oriented towards the ligands) is heightened by increased repulsion due to shortened e_g orbital-ligand distance during pressure-induced unit cell distortion, resulting in an overall larger CFSE. When the CFSE surpasses the spin-pairing energy, the spin-pairing transition of 3d electrons is more favorable than jumping the energy gap to achieve aligned spins. For example, the low-spin state with all six 3d electrons paired (S=0) in Fe²⁺ occurs at high pressures in the lower-mantle ferropericlase

spectroscopy (SMS), because these results act as site-specific fingerprints, assisting in the assignment of the valence and spin states of iron in the lower-mantle minerals.

2 Electronic spin transition of Fe^{2+} in Fp

Ferropericlase has a cubic rock-salt structure and forms a solid solution between periclase (MgO) and wüstite (FeO). A high-spin (HS) to low-spin (LS) transition of Fe²⁺ in the octahedral site occurs when the crystal field splitting energy exceeds the Hund's-rule exchange energy, causing the material to become low-spin and diamagnetic [1–4]. The measured values of QS, \sim 0.8 mm/s, and CS, \sim 1.2 mm/s, from Mössbauer spectra of Fp under ambient conditions are consistent with predominant high-spin Fe²⁺ in the octahedral coordination (Fig. 3) [5]. The concentration of the high-spin Fe³⁺ in ferropericlase, however, is typically very low (e.g., under the detection limit of the MS technique). At room temperature, the QS of the high-spin Fe²⁺ increases with pressure up to \sim 30 GPa, plateaus until \sim 60 GPa, and disappears with further pressure increase [5]. The CS of the high-spin state decreases with pressure increase; a noticeable drop of the CS occurs at approximately 60 GPa. The simultaneous disappearance of the QS and the drop of the CS at approximately 60 GPa are consistent with a spin transition of iron in Fp [5]. The ratio of the high-spin to low-spin states of iron in Fp, as a function of pressure, can be derived from the



modeling of the SMS spectra with changes in the QS and CS values. Depending on experimental conditions, a narrower to broader width of the spin transition has been reported [1, 5]. Although (Mg,Fe)O has always been regarded as a fully disordered solid solution system without noticeable short-range order at ambient conditions, recent Mössbauer analyses have demonstrated broadening of the high-spin Fe²⁺ quantum beats associated with significant changes in the short-range order, which deviates from the behavior of the homogeneous solid solution in Fp. These studies have indicated that the Fe²⁺ ions can form localized clusters in nearest-neighboring environments under high pressures. The short-range ordering of Fe²⁺ ions can cause an extension of the width of the spin crossover in (Mg,Fe)O at high pressures.

The electron shell of the high-spin Fe²⁺ ions in the octahedral coordination is spherically asymmetric with S=2 and $t_{2g\uparrow}^3e_{g\uparrow}^2t_{2g\downarrow}^1$, whereas Fe²⁺ ions in the low-spin state are more spherically symmetric with S=0 and $t_{2g\uparrow}^3t_{2g\downarrow}^3$ (the arrows indicate spin up or spin down whereas the superscripted numbers represent the number of the occupied electrons). The spin crossover arises from the condition in which the thermal energy at high P-T is sufficient to overcome the energy difference between the high-spin state and low-spin state [4]. Therefore, the local 3d electronic structures of the iron ions in Fp are strongly temperature dependent [4]. Most importantly, recent results show that the electronic spin-pairing transition of iron in Fp occurs over a very narrow range of pressure at room temperature but turns to a spin crossover with an extended transition pressure at the lower-mantle temperatures [4].

3 Spin and valence states of iron in Pv

Perovskite, with 5–10 mol% of Fe and Al, is the most abundant silicate mineral in the lower mantle, and exists at 660 km depth to several hundred kilometers above the core-mantle boundary. Iron in Pv exists in both Fe^{2+} and Fe^{3+} states and can occupy one of two crystallographic sites, the large dodecahedral Mg site (the A site) and the small octahedral Si site (the B site) [6–11] (Fig. 2). Current consensus on the site occupancy is that Fe^{2+} mainly occupies the A site whereas Fe^{3+} can occupy both the A and B sites [6–11]. Several studies have further shown that Fe^{2+} may self-disproportionate into Fe metal and Fe^{3+} , creating Fe^{3+} -enriched Pv. That is, the significant amount of Fe^{3+} in Pv is not a result of the oxidation state of the lower mantle, which is expected to be relatively reducing, but due to the crystal chemistry of Pv at high pressure. The Fe^{3+} content in Pv is also found to vary with the amount of the Al^{3+} likely through charge-coupled substitution with the Si^{4+} in the octahedral site [4, 6]. The Fe^{3+} content in Pv is approximately 20% without the presence of Al and 50–75% with the presence of Al^{3+} .

The reported spin transitions in Pv are much more complex than those in Fp, likely due to the low-symmetry oxygen ligand field and multiple site occupancies. These studies have shown that both Fe^{2+} and Fe^{3+} exist in the high-spin state in both of the A and B sites under ambient conditions [6–11]. Mössbauer analyses show that the high-spin Fe^{2+} exhibits a QS value of \sim 1.6 to 2.2 mm/s and a CS of \sim 1 mm/s in the A site. Interpretations of the high-pressure Mössbauer analyses have been quite different, although their main results on the QS values were similar—both experimental and theoretical studies have reported extremely high QS values



of Fe²⁺ (as high as \sim 4.4 mm/s) at high pressures [6–11]. The relative area of the high-QS doublet (3.4-4.4 mm/s) increases with pressure at the expense of the low-QS doublet (1.6–2.2 mm/s), indicating that the transition involves Fe²⁺ in the A site [7, 8]. The high-OS doublet becomes the dominant spectral feature in the high-pressure Mössbauer spectra at above approximately 30 GPa. Particularly, the new quadrupole doublet with the extremely high QS appears to have a very narrow linewidth and high CS (\sim 1.1 mm/s) [7]. Together with X-ray emission spectroscopic (XES) analyses for the total spin momentum of iron in Pv, the occurrence of the Fe²⁺ site with the extremely high QS, very narrow linewidth, and relatively high CS has been assigned to be an intermediate-spin Fe^{2+} with a total spin momentum of one (S=1) in the A site [7]. That is, a high-spin to an intermediate-spin crossover occurs in Pv at around 30 GPa, and Fe²⁺ remains stable in the intermediate-spin state at above 30 GPa in the lower mantle. At higher pressures, the intensity of the high QS component in the SMS spectra of PPv decreases, whereas the intensity of a new component with very low QS of less than 0.5 mm/s and a CS of ~0 mm/s increases. This new component was assigned to the low-spin Fe²⁺ occurring at 120 GPa and high temperatures.

However, it has been realized that the occurrence of the intermediate-spin state is very rare in geological materials and that high QS in iron does not necessarily imply the occurrence of the intermediate-spin state. For example, iron in almandine garnet is in the high-spin state with a QS of \sim 3.5 mm/s under ambient conditions. Theoretical calculations have been performed to interpret the spin states and the extremely high QS values of Fe²⁺ in Pv. Although not always in agreement with each other, these calculations show that the intermediate-spin state is not stable over all lower-mantle pressures, irrespective of the exchange-correlation functions used in the calculations [9]. The high-spin Fe²⁺ with QS of 2.3–2.5 mm/s is more stable at relatively low pressures, while the high-spin Fe²⁺ with QS of 3.3–3.6 mm/s is more favorable at higher pressure [9]. Based on these calculations, the extremely high QS site is a result of the iron atomic-site change, in which Fe ions move away from the central positions in the A site, rather than a spin-pairing transition.

For Fe³⁺ in Pv, it has been shown both experimentally and theoretically that Fe³⁺ enters into both A and B sites, suggesting a charge-coupled substitution mechanism [8, 10]. High-spin Fe³⁺ is known to exhibit very low QS compared to Fe²⁺ because all five 3d electrons are unpaired and form relatively spherical orbitals. Theoretical calculations further suggest that the Fe³⁺ in the A site has an even lower QS value than the B site Fe³⁺ [10]. Combined SMS and XES results for Fe³⁺-containing Pv suggest that the low-spin population in the B site gradually increases to 100% with pressure up to 50–60 GPa, whereas Fe³⁺ in the A site remains high-spin to at least 136 GPa, consistent with recent theoretical calculations [8, 10]. SMS analyses further suggest that A site Fe³⁺ remains in the high-spin state in Al-bearing Pv, in which Fe³⁺ is expected to predominantly exist in the A site [8]. Therefore, these results indicate that the Fe³⁺ in the octahedral B site undergoes a high-spin to low-spin transition in Pv.

The QS of the Fe^{2+} in the A site and the Fe^{3+} in the B site exhibit exactly the opposite trends across the high-spin to the low-spin transition, and can be understood through their orbital occupancies. The low-spin Fe^{2+} in the A site located near the center of the site forms a 3d charge density with a cubic-like shape that barely contributes to the effective electric field gradient, leading to a very small QS [8, 10]. The high-spin Fe^{3+} also has a small QS (a small effective electric field gradient) in the



A and B sites because of an almost spherically-shaped electron charge distribution [8, 10]. However, the 3d orbitals of the low-spin Fe³⁺ contribute more significantly to the effective electric field gradients and lead to a larger QS.

4 Spin and valence states of iron in PPv

Post-perovskite is expected to be the most abundant phase in the Earth's core-mantle boundary region. Deciphering the spin and valence states of iron in PPv at relevant P-T conditions provides new insight into the properties of the region, including seismic discontinuities, rheology and plasticity, dynamic evolution and formation of superplumes, thermal gradients, core-mantle heat flux, and chemistry [11]. The PPv phase is found to be stable in the CaIrO₃-type structure (*Cmcm*). However, a number of kinked structures, formed by sliding the {010} planes of the Pv structure with variation in the stacking sequence of SiO₆ octahedral layers, have also been reported. Thus far, the presence of the kinked PPv phase has been found to increase the amount of Fe³⁺ in the sample. Experimental SMS results show Fe²⁺ likely exists in the bipolar-prismatic site and exhibits extremely high QS of 3.8-4.5 mm/s and relatively high CS in PPv [11]. Complementary high P-T XES studies, which are sensitive to the total spin momentum of iron, suggest that iron predominantly exists in the intermediate-spin Fe^{2+} state with S=1 in the CaIrO₃-typed PPv at relevant P-T conditions of the lowermost mantle [11]. Since the intermediate-spin Fe²⁺ in PPv is found to be stable at 25 mol% and 40 mol% Fe, it is conceivable that the intermediate-spin Fe²⁺ is stable over a wide range of Fe content in PPv relevant to the D" region, where Fe-enrichment may be expected [11]. Similar to the discussion for Fe²⁺ in Pv, however, theoretical calculations suggest that Fe²⁺ in PPv is in the high-spin state at all mantle pressures, although theoretical QS values of the Fe²⁺ and Fe³⁺ remain to be computed.

SMS results on Fe³⁺-rich PPv show that Fe³⁺ exists in two different sites, one site with a high QS of \sim 2 mm/s and another with a low QS of 0.3 mm/s, similar to the Fe³⁺ in Pv [12]. The low QS site is assigned to be the high-spin Fe³⁺ in the bipolar-prismatic site, whereas the high QS site with QS of \sim 2 mm/s is assigned as the low-spin Fe³⁺ in the octahedral site [12]. Based on the hyperfine parameters associated with the site assignments, the Fe³⁺ in the octahedral site undergoes a high-spin to low-spin transition at high pressures, causing the QS to increase from the high-spin state to the low-spin state [12]. In this mixed spin-state scenario, Fe³⁺ enters the site through charge-coupled substitution, as also suggested by theoretical calculations. Because the ionic radius of the low-spin Fe³⁺ is smaller than that of the high-spin Fe³⁺, the low-spin Fe³⁺ is more stable in the smaller octahedral site of PPv.

The notable difference between the Mössbauer parameters of the Pv and PPv phases is that the QS of the low-spin Fe^{3+} site is much smaller in PPv. The smaller QS of the low-spin site in PPv can be explained by a less distorted octahedral site, since distortion from a cubic environment around the iron nucleus results in an increased QS value. Since PPv is the high-pressure polymorph of Pv, the QS of Fe^{2+} in PPv could be similar or related to that of Fe^{2+} in Pv as both phases exhibit extremely high QS values of around 4 mm/s. The calculated QS of Fe^{2+} in Pv is 3.3–3.6 mm/s at lower mantle pressures, consistent with the QS values for Fe^{2+} from SMS analyses in PPv [7–12].



5 Conclusions

The disappearance of the QS and drop in the CS of the octahedrally-coordinated Fe²⁺ in ferropericlase indicates the high-spin to low-spin transition at high pressures. The spin transition of iron in ferropericlase turns into a wide spin crossover at lowermantle temperatures. Iron exists in the Fe²⁺ and Fe³⁺ states in both perovskite and post-perovskite. Both Fe²⁺ and Fe³⁺ exist in the high-spin state in perovskite under ambient conditions. Most recent studies have observed extremely high QS values of the Fe²⁺, as high as \sim 4.4 mm/s, in perovskite under high pressures. The relative area of the high-OS doublet increases with pressure at the expense of the low-OS doublet, and has been assigned as an intermediate-spin Fe²⁺ in the A site, occurring at approximately 30 GPa. However, recent theoretical calculations support the notion that the extremely QS site is a result of the atomic-site change rather than a highspin to intermediate-spin transition. At higher pressures, a new doublet component was assigned to the low-spin Fe²⁺ occurring at 120 GPa and high temperatures. Recent studies on Fe³⁺-containing perovskite suggest that Fe³⁺ in the octahedral B site undergoes a spin-pairing transition in perovskite, whereas Fe³⁺ in the A site remains high-spin to at least 136 GPa.

Fe²⁺ likely exists in the bipolar-prismatic site with an extremely high QS of 3.8–4.5 mm/s in post-perovskite, which has been assigned to the intermediate-spin Fe²⁺ state with S=1. However, theoretical calculations have found the intermediate-spin state unstable at lower-mantle pressures. Fe³⁺ exists in two different sites, the high-spin Fe³⁺ in the bipolar-prismatic site and the low-spin Fe³⁺ in the octahedral site. These site assignments indicate that the Fe³⁺ in the octahedral site undergoes a high-spin to low-spin transition at high pressures through charge-coupled substitution.

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